EFFECT OF RUPTURE IN A FISSION GAS HOLDUP BED*

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Abstract

A gas adsorption bed operated under pressure will lose a portion of the adsorbed gases in the event of an accidental rupture of the container and release of pressure. Theoretical and laboratory studies show that the fractional release of fission gases will be smaller than the ratio of the initial to the final bed pressure and that the fraction of krypton released will be greater than the fraction of xenon released under the same conditions.

I. Introduction

Currently the short lived isotopes of krypton and xenon constitute the bulk of the radioactivity released to the atmosphere from nuclear reactors. The release of gases contaminated with these radionuclides may be delayed by a holdup system, and if the holdup time is sufficiently long (e.g., 120 days), the off gas stream may be effectively decontaminated of all short lived fission gas nuclides. The ten year nuclide, 85Kr, cannot be controlled by such a holdup system, but fortunately the fission yield of this nuclide is small. A major problem in the design of a fission gas holdup system is the space required for an effective system. required storage volume may be reduced considerably either by filling the holdup tanks with an adsorbent material (such as charcoal) and/or by operating the system under pressure. Operation under pressure, however, presents the potential situation in which the bed develops an accidental leak with total loss of pressure and accompanying release of a portion of the adsorbed gases. This paper presents an evaluation of this type of accident taking place in a pressurized bed filled with adsorbent. The study was performed by the author in conjunction with the design of a fission gas holdup system for the Fast Flux Test Facility.

II. Accident Considerations

Several types of releases are possible in the case of the development of a leak in a pressurized bed. But a violent rupture

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in which the bed container essentially disintegrates with portions of the adsorbent ejected appears highly unlikely because the maximum operational pressure generally considered for such a bed would be restricted to eleven atmospheres (absolute) or lower. A more probably accident situation is a slow leak, and in this case, the most important points to consider are:

- a. In the presence of an adsorbent, the fission gases, krypton and xenon, are much less mobile than their carrier gas (which for the FFTF off gases will be argon). For this reason, it would appear that the fraction of fission gases escaping from the bed will be significantly smaller than the fraction of carrier gas.
- b. As a result of radiodecay at steady state, the concentration of fission gas will be highest near the bed inlet and lowest near the bed outlet. If a leak develops near the feed input, the possibility exists that only a small release of carrier gas could sweep out an appreciable fraction of the fission gases contained within the bed.

The net effect of these two opposing factors on fission gas leakage can be calculated on the basis of the analysis developed below.

III. Development of Theory

To establish a model for numerical analysis, we assume that there is a rupture at or near the entrance to the bed diagrammed in Figure 1. As was pointed out above, this would be the point of highest fission gas concentration within the bed and would, therefore, be the point where maximum release would occur.

The fission gas adsorbed in the bed nearest the point of rupture will be swept out of the bed by the escaping gases, whereas the fission gas adsorbed deepest in the bed will be retained. The first example considered here is a bed in which a leak develops at the bed inlet (the point x=0 in Figure 1). The analytical procedure used here is to find the point, z, in the bed dividing the fission gas which escapes from that which is retained. The following assumptions have been made in locating the point, z.

- a. the release occurs under isothermal conditions;
- b. the fission gases are highly diluted by the carrier gas;
- c. both the fission gas and the carrier gas follow linear adsorption isotherms; and
- d. the pressure release is confined to a single bed.

Under these conditions, the instantaneous flow of all gases at any point in the bed is proportional to the distance from the point, L, which represents the effluent end of the bed. Also, the instantaneous flow velocity of a fission gas is equal to the instantaneous flow velocity of the carrier gas divided by the ratio

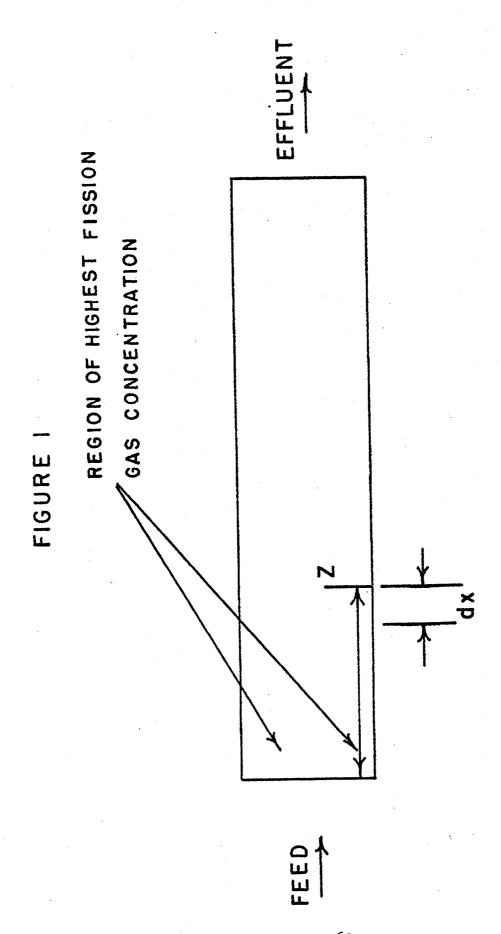


DIAGRAM OF FISSION GAS ADSORPTION BED

of their adsorption coefficients. It follows directly that the differential equation giving the distance moved by the carrier gas is:

$$-\frac{d(L-x)}{L-x} = \frac{dP}{P}, \tag{1}$$

where P = absolute pressure in the bed, and further the incremental distance moved by a fission gas is given by:

$$-\frac{d(L-x)}{L-x} = \frac{dP}{aP} \tag{2}$$

where $a = ratio \begin{cases} \frac{adsorption\ coefficient\ for\ fission\ gas\ charcoal}{adsorption\ coefficient\ for\ carrier\ gas\ charcoal} \end{cases}$

On integration, Equation (2) becomes

$$z^{0} - \frac{d(L-x)}{L-x} = \int_{P_{1}}^{P_{atm}} \frac{dP}{aP}$$
 (3)

where P_{atm} = atmospheric pressure

P; = initial pressure.

Then

$$\ln(\frac{L-z}{L}) = \frac{1}{a} \ln (P_{atm}/P_i)$$
 (4)

or

$$z = L \left\{ 1 - \left(\frac{P_{atm}}{P_i} \right)^{1/a} \right\}$$
 (5)

During normal operation, the radioactive fission gases will have an exponential decrease in activity as a result of natural decay while passing through the bed. Therefore, it can be assumed that the distribution of fission gas activity as a function of distance from the entrance can be expressed as: $g(x) = be^{-\alpha x}$.

Whatever the function g(x) is, the fraction, f, of a fission gas nuclide released from the bed will be

$$f = \int_{0}^{z} g(x)dx / \int_{0}^{L} g(x)dx$$
 (6)

When $g(x) = be^{-\alpha x}$,

$$f = \frac{1 - e^{-\alpha Z}}{1 - e^{-\alpha L}} \tag{7}$$

or substituting the value for z,

$$f = \frac{1 - e^{-\alpha L} \left\{ 1 - \left(\frac{P_{atm}}{P_{i}} \right)^{1/a} \right\}}{1 - e^{-\alpha L}}$$
(8)

When $\alpha L << 1$, only a small fraction of fission gas undergoes decay while passing through the bed and a nearly even initial distribution of fission gas across the bed results. For this condition, the equation for f reduces to

$$f = 1 - \left\{ \frac{P_{atm}}{P_{i}} \right\}$$
 (9)

which is essentially the equation derived by Jury (1) for a release from an adsorption bed having a uniform initial distribution of dilute fission gas.

For the case of ideal mass transfer (2)

$$\alpha = \frac{\lambda \rho K_{o}(fission gas)}{V_{s}}$$
 (10)

and

$$a = \frac{K_o(fission gas)}{K_o(carrier gas)}$$
 (11)

where

 ρ = density of the adsorbent, gms/cm³

 $K_0 = adsorption coefficient, cm³/gm$

V_s = superficial carrier gas velocity, cm/sec

 λ = isotopic decay constant, sec⁻¹

The bulk adsorption coefficient, K, can be defined in many ways: to avoid confusion, K is defined here as the volume of a gas in a gram of bulk adsorbent measured at the partial pressure of the gas in the adsorbent. With this definition and the previous assumption of a linear isotherm, the bulk adsorption coefficient, K, becomes independent of the partial pressure of the adsorbed gas. The superficial velocity, V, is assumed to be measured at the pressure inside the bed before the rupture occurred.

IV. Experimental Results

The theory presented here was tested on a laboratory scale by measuring the fractions of $^85\mathrm{Kr}$ and $^{133}\mathrm{Xe}$ released from a charcoal bed following a sudden loss of pressure. The experimental procedure was to pass, under pressure at room temperature, argon plus trace quantities of either $^{85}\mathrm{Kr}$ or $^{133}\mathrm{Xe}$, through a charcoal bed until the effluent concentration equaled the input concentration. At this point the bed was saturated with the fission gas, and the bed was then isolated and depressurized. The adsorption coefficient for argon could be determined from the volume of argon released from the bed. The activity of the fission gas in the released argon was also measured. By sweeping the bed with uncontaminated argon, the residual activity of the fission gas in the bed was also found, and from the total amount of fission gas in the bed, at saturation, the adsorption coefficient for the fission gas was also calculated. Thus, during a single run, both fractional release of fission gas, as well as all the parameters needed to calculate the theoretical fission gas release, were measured. Table 1 gives the experimental and theoretical fission gas releases for $^{85}\mathrm{Kr}$ and $^{133}\mathrm{Xe}$ from beds pressurized to 2 and to 10 atmospheres (gage).

As predicted from theory, the fractional release of fission gases was smaller than the ratio of the initial to the final bed pressure. Also, as predicted, the fraction released increased with initial pressure, and under similar conditions, the fraction of krypton released was significantly greater than the fraction of xenon released.

It is important to note that the observed releases were some-what less than those calculated from the assumption of an isothermal release. The fact is that the desorption of large volumes of the argon carrier gas resulted in a simultaneous cooling of the beds during the depressurization, and this cooling in effect changed the value of the parameter "a" during the release. The analysis could

TABLE 1

FRACTIONAL RELEASE OF FISSION GASES FOLLOWING RAPID DEPRESSURIZATION

Carrier Gas: Argon; Temperature: Ambient

Predicted Fractional Release		80	σ.	\O	91	[]	8 1	9+
Pred Fraction	0.21	0.18	0.48	94.0	0.016	0.011	0.048	0.046
Observed Fractional Release	0.16	0.14	0.32	0.34	0.011	0.008	0.018	0.017
Initial Pressure (gage)	30 psi	30 psi	150 psi	150 psi	30 psi	30 psi	150 psi	150 psi
Weight of Charcoal in Bed	480g	480g	480g	480g	480g	480g	144g	144g
Fission	85 _{Kr}	$^{85}{ m Kr}$	$^{85}_{ m Kr}$	85 _{Kr}	133Xe	133 <u>x</u> e	$^{133}_{ ilde{ imes}_{ m Ke}}$	133 xe
Run	П	7	\sim	4	7	9	7	œ

be altered to take this effect into account, but such a procedure would then not be valid for a slow, isothermal release. For this reason we propose that the simple formulas developed here be retained with the understanding that for sudden releases they may be somewhat conservative.

V. Sample Calculations

To demonstrate typical fission gas release values based upon the theory developed above, sample calculations were performed for the situation assumed to exist in a bed pressurized to 10 atmospheres (gage).

In a set of sample data for operation of a holdup bed at ambient temperature (25°C)

$$\rho = 0.48 \text{ gm/cm}^3$$

$$K_{O} = 357 \text{ cm}^3/\text{gm for Xe}$$

=
$$28 \text{ cm}^3/\text{gm for Kr}$$

=
$$6.64 \text{ cm}^3/\text{gm for Ar}$$

$$V_s = 0.25 \text{ cm/sec}$$

L = 300 cm

P = 10 atmospheres (gage)

Table 2 gives the curies/sec, q, for each radionuclide in the feed to the bed. This rate of release is calculated from an FFTF Deisgn assuming four 2 MW closed loops and 1% defected fuel in reactor. From Equation (12) the activity of each nuclide in the holdup bed can be calculated.

$$q = \frac{\dot{q}}{\lambda} (1 - e^{-\alpha L}) \tag{12}$$

where q equals the curies of each nuclide contained in the bed. From Equation (8) the fractional release of each nuclide is calculated, and this as well as the total release is also given in Table 2. By an analogous procedure the fission gas release from a leak in the middle or at the outlet of the bed was calculated and the results are also given in Table 2.

VI. Commentary and Conclusions

As may be noted from Table 2, the development of a leak in a fission gas holdup bed, pressurized to 10 atm (gage) and operating under equilibrium conditions, will results in a greater fractional release of the krypton isotopes. This is as would be expected because of the greater mobility of this gas. Further, if the leak occurs near the bed input, there is a greater fractional release

TABLE 2

CALCULATED RELEASES FROM A FISSION GAS HOLDUP BED

Initial Pressure: 10 atm (gage)

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Effluent End	Curies	12.63	173.69	5183.61	531.08	527.99	1832.20	0.21	916.16	2829.60
Eff1ue	Fraction	0.040	0.030	0.037	0.002	0.246	0.349	0.428	0.194	0.304
of Bed	Curies	13.49	244.56	6013.14	4146.54	847.01	2236.86	0.20	1850.02	3872.34
Middle	Fraction Released	0.043	0.042	0.043	0.022	0.395	0.426	0.423	0.367	0.416
Input End	Curies	14.41	344.35	6975.68	32375.20	1358.79	2730.91	0.21	3506.15	5299.35
InduI	Fraction	0.046	090.0	0.050	0.173	0.634	0.520	0.432	769.0	0.569
	Curies in Bed at Steady State	309.5	5726.8	138408.4	186818.3	2140.7	5244.5	0.5	5027.9	9302.5
	Input Curies/Day	139.2	3369.0	67650.0	342000.0	23586.0	39150.0	5.6	67920.0	82440.0
	Nuclide 131m	ә <u>х</u> штст	$133m_{\mathrm{Xe}}$	133_{Xe}	$_{\rm 3.135_{Xe}}$	83m _{Kr}	$85 \mathrm{m_{Kr}}$	$^{85}\mathrm{Kr}$	$^{37}\mathrm{Kr}$	$88_{ m Kr}$

of shorter lived radionuclides. Since these nuclides constitute the bulk of the radioactivity near the area of the assumed leak, this situation is as would be expected.

The overall release of fission gases is calculated to be 14.9 per cent of the total radioactivity contained within the bed if the release occurs near the bed input. Similar calculations for assumed leaks in the middle and near the outlet end of the bed showed radionuclide releases only of 5.4 and 3.4 percent, respectively of the total radioactivity contained within the bed.

Furthermore, if the adsorption bed were to be operated at cryogenic temperatures, the ratio of adsorption coefficients (the factor"a") would increase substantially. Under these conditions, the fraction of fission gases released from a bed could be significantly smaller than those calculated here. In addition, we have assumed an initial pressure of 10 atmospheres (gage). For operation at lower pressures, smaller fractional releases on depressurization would be expected.

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DISCUSSION

JONAS: In order to integrate the right side of your equation you would have to assume that "a" was not a function of the pressure changes?

UNDERHILL: This would be the case for linear isotherms.

JONAS: It would also indicate that the isotherms of the fission gas and the carrier gas were approximately the same. I wonder if that was one of the causes for the variance between the observed and calculated data.

UNDERHILL: We know from the results of Förster and some other work we did, that there is a change in the adsorption coefficient of krypton at high pressures. I think there is probably some effect of this in the results. But, again, there was a very, very strong cooling effect and I think that was the primary reason for the deviation between theory and experiment.

JONAS: Yes; I agree. I guess what I was pushing for is the question of whether the cooling effect is adding to or subtracting from the deviations.

UNDERHILL: I see your question, and it's really an unanswerable one, unless we do further research.

WYMER: You talk about rapid depressurization. I wonder how rapid; and how did you do it experimentally?

UNDERHILL: We simply opened up a valve, releasing everything through a quarter-inch copper tubing.

WYMER: How long did it take?

UNDERHILL: There was an initial swoosh of material and then we waited about two hours to allow the temperature of the bed to become the ambient temperature.

WYMER: You mean a one-second swoosh or a two-second swoosh?

UNDERHILL: It's hard to say.

MICHELS: Have you considered the case in which the bed ruptures and you have a gas stream flowing over the charcoal?

UNDERHILL: Probably we could have, but we haven't considered that case up to now.

MASS AND HEAT TRANSFER OF KRYPTON-XENON ADSORPTION ON ACTIVATED CARBON

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Abstract

A large number of boiling water reactors are being equipped with ambient temperature-carbon adsorption off-gas systems. The selection of adsorbent for these systems is often based on limited experimental data resulting in non-optimized processes. The data presented here was generated to assist in the selection of the adsorbent and operating conditions for the off-gas systems utilizing the ambient or near ambient adsorption process.

I. Carbon Optimization.

The adsorption of krypton and xenon on various adsorbents has been extensively studied in the past. Most investigators have used only one adsorbent, or adsorbents of very widely varying properties; therefore, the resulting data did not permit optimization of a single base adsorbent. (1)

Although Lad and Young reported that lower surface area identical base adsorbents exhibited higher capacity toward xenon at low concentrations than high surface area carbons and Mahajan and Walker reported similar results for krypton adsorption on activated carbons, many investigators selected, and most results were reported on, high surface area adsorbents. (2) (3)

The experiments to evaluate optimum surface area were performed on steam activated coconut base carbons, subjected to various activation levels. The carbons used had surface areas of $525 \pm 50 \text{ m}^2/\text{g}$,

$$680 \pm 50 \text{ m}^2/\text{g},$$

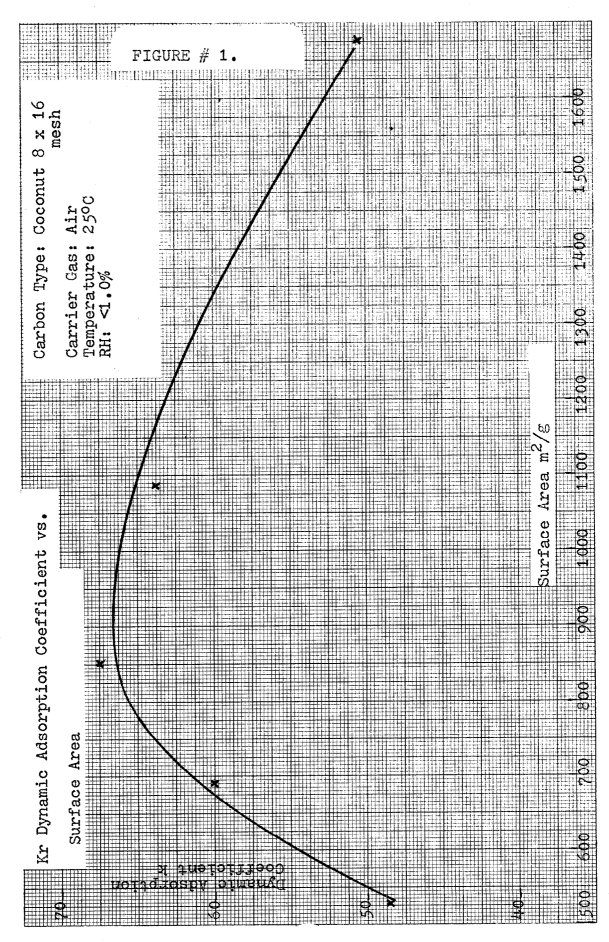
$$850 \pm 50 \text{ m}^2/\text{g}$$

$$1085 \pm 50 \text{ m}^2/\text{g}$$

using the BET nitrogen adsorption method.

The dynamic adsorption coefficients were determined using Kr-85 tracer at 0.001 mmHg pressure in air. The column dimensions were 6.0 inch ID and 3 foot bed depth. The temperature was 25°C, the superficial velocity 2 fpm and the relative humidity was kept below 1.0%. The carbon adsorbents used were equilibrated with tracer free air under the experimental conditions. An Overhoff & Associates Model 120 ion chamber was used to monitor Kr-85 breakthrough.

The obtained experimental data is shown on Figure No. 1. The optimum surface area of this adsorbent series is near 800 m²/g for krypton adsorption delay. The probable reason for the observed phenomenon is that during the activation process not only the surface area increases but the pore size distribution of the activated carbon also increases. During the first phase of the activation process new pores are developed while in successive stages of the activation the pore size is enlarged. Due to the fact that particularly at low concentration adsorption takes place preferentially in the smallest diameter pores, the use of the lower surface area adsorbent is advantageous if the pore diameters



are small. Mahajan (3) evaluated carbonaceous molecular sieves also showing that if by unique processing the pore diameter can be kept small. even at higher surface areas, a superior carbon can be developed for krypton adsorption. However. the preparation of carbonaceous molecular sieves at the present time is not an economically competitive process. Thus based on the experimental results, the optimum pore diameter for krypton adsorption, in steam activated coconut carbons. is developed at approximately 800 m²/g and further increase in surface area does not result in increased delay times due to the enlargement of the pores. Literature data (4) does indicate that under cryogenic conditions this relationship between pore diameter and krypton, xenon delay times is not valid. The temperature dependence of this relationship is being evaluated at the present time.

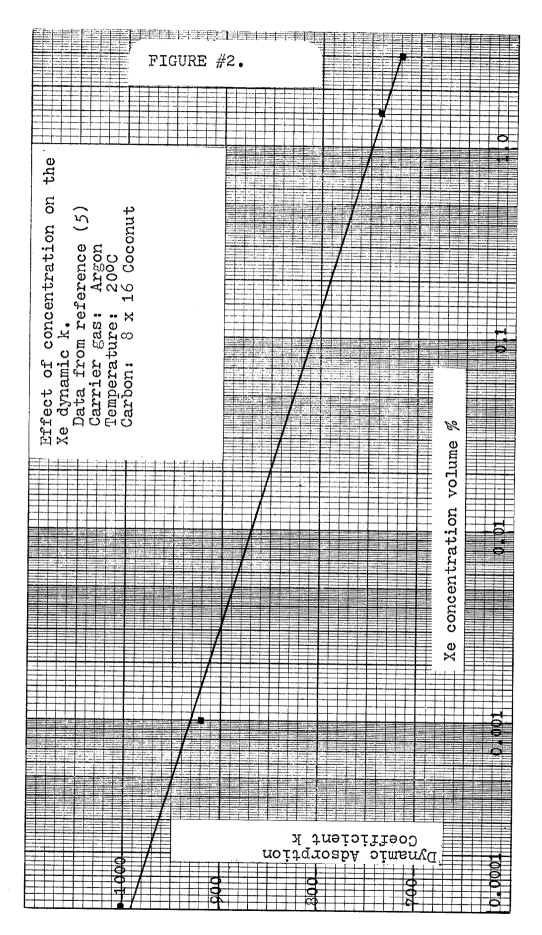
Even more significant differences are observed if the Kr or Xe dynamic k is based on adsorbent volume rather than weight. The surface area of activated carbons is inversely proportional to their apparent density; therefore the volumetric Kr-Xe adsorption capacity shows even more advantage for using the higher density, lower surface area coconut carbons.

The position of the krypton and xenon adsorption isotherms determined under static conditions also changes relative to each other on various pore diameter and surface area carbons. Therefore it is important that any single concentration point evaluation is performed at or as close to the actual concentrations as possible. To demonstrate this concentration dependence, literature data plot obtained under various concentrations on the identical adsorbent is shown on Figure No. 2. The data was obtained under dynamic conditions from argon carrier gas. (5)

The data indicates that contrary to other data the slope of the adsorption isotherm, i.e. both the static and dynamic k, changes even at the very low noble gas concentrations.

II. Dynamic Considerations.

As was shown in an earlier report (6), the movement of a noble gas molecule through the adsorbent bed is also dependent on the slope of the adsorption isotherm, i.e. the dynamic (or static) adsorption coefficient k.



The rate of movement of the adsorption mass transfer zone in case of fixed superficial velocities is a function of the distribution of the adsorbate (in this case Kr and Xe) between the carbon and the gas phase.

$$u = v \frac{1}{a/P_0 \epsilon} .$$
 [1]

The proportionality factor $a/P_0\epsilon$, giving the distribution of the Kr and Xe between the adsorbed and the gas phase, can be substituted by the adsorption coefficient k if the adsorbate quantity in the void volume is negligible, resulting in

$$u \approx q \frac{1}{k}$$
 [2]

Thus the rate of movement of the adsorption mass transfer zone, i.e. the rate of movement of the Kr or Xe molecules through the adsorbent bed, will be accelerated by increasing the superficial velocity and will be decreasing in proportion to the adsorption coefficient.

If the adsorption isotherms of the particular adsorbateadsorbent system are nonlinear over wide ranges, the nearest available data range k value can only be substituted.

At low gas velocities, particularly when using ion chambers which may dilute adsorbate concentrations, twin columns of two different bed lengths should be used. The obtained results

$$u = \frac{L_2 - L_1}{t_2 - t_1} \approx q \frac{1}{1 + k}$$
 [3]

will give more precise approximation of the rate of mass transfer zone movement.

Under very low velocity conditions, resulting in u of less than 3 cm/min Kr and Xe mass transfer zone movement in activated carbon beds, the longitudonal diffusion in the adsorber considerably increases the length of the mass transfer zone. Similar results were reported by Illes for CO2, C2H2, C2H6 and C3H8 adsorption on activated carbon. (7)

III. Heat Transfer in Carbon Beds.

Although numerous correlations exist for packed granular bed heat transfer data, the mass velocity in these correlations was obtained at high Reynolds numbers. A theoretical derivation was developed to show the heat transfer coefficient versus gas velocity dependency through a wide velocity range. (8)

The equation representing the derivation

$$Nu = 0.2 + 0.03Pr^{0.33}Re^{0.54} + 0.35Pr^{0.356}Re^{0.58}$$
 [4]

was evaluated in a 6.0 inch ID 3 foot long jacketed vessel. Figure No. 3 shows the curve corresponding to equation [4] and the data points obtained experimentally on an 8 x 16 mesh carbon in the 400-100°C range in nitrogen. The following parameters were used or measured.

Average particle size: 0.128 cm

Average particle surface: 0.10 cm²

Average particle volume: 0.002050 cm³

Average number of particles: 278

Bed void fraction: 0.42

Surface area external: 27.0 cm⁻¹

Sphericity: 0.719

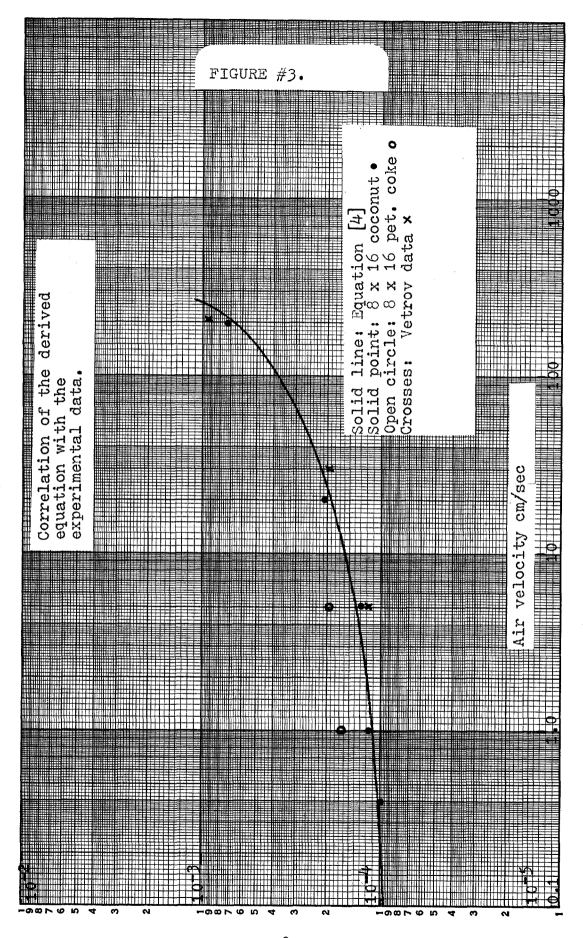
Carbon thermal conductivity: 3.10×10^{-4}

cal/(cm)(sec)(°C) at 100°C

Carbon heat capacity: 0.20 cal/(g)(°C) at 100°C

Prandtl No.: 0.659

Reynolds No.: 0.777v



The values reported by Vetrov (9) for chips and steel shot were also plotted and show close agreement with the obtained data. The developed equation [4] agrees in order of magnitude with the conventional (Wilke-Hougen, Gamson-Hougen, Colburn-Chilton, etc.) equations only in the turbulent flow and very low velocity ranges. The values observed exclude any oxidation effects which may occur at high temperatures (above 120°C) in oxygen or oxidizing agent containing carbon beds.

Equation [4] confirmed by experimental data, results in a shape which points out the nonlinearity of the heat ransfer coefficient-air velocity relationship over the turbulent, intermediate, laminar flow and "chimney effect" velocity ranges.

Alimited number of experiments were performed using near spherical carbon particles of similar size and properties as the average size of the coconut shell carbon used. The data resulted in slightly higher α values than those obtained with the prismatic shape coconut carbon even when the latter is subjected to sphericity correction.

Whether this phenomenon is caused by the variation of the angle of attack or by the packing density difference in the non-uniform size coconut carbon beds is not clear at this time.

Summary.

The influence of carbon adsorbent structural properties were evaluated for noble gas removal systems. The effect of the obtained adsorption coefficients is discussed in relation to the movement of the mass transfer zone through the adsorbent bed. A heat transfer equation is presented for heat transfer on packed carbon beds; the equation is substantiated by measured data points and literature values.

Symbols.

- a = volume of adsorbate adsorbed in 1 cm³ adsorbent cm³/cm³.
- k = adsorption coefficient, volumetric cm³/cm³.
- L = adsorbent bed length cm.
- P_0 = adsorbate concentration fraction at inlet cm³/cm³.
- q = true velocity through the adsorbent bed cm/min.
- t = time elapsed to breakthrough minutes.
- u = rate of movement of the adsorption wave
 front cm/min.
- v = superficial velocity of the carrier gas cm/min.
- α = film heat transfer coefficient cal/(cm)²(sec)(°C).
- ϵ = void fraction of adsorbent bed cm³/cm³.

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DECONTAMINATION OF HTGR REPROCESSING OFF-GASES

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Abstract

The off-gas from the reprocessing of HTGR fuel is unique among nuclear fuel reprocessing off-gases because of the large amount of CO2 produced by the burning of the fuel element. An HTGR fuel element contains about ten times as much graphite as heavy metals. The relatively high temperature of the burning process (0800°C) releases virtually all of the volatile fission products, as well as troublesome amounts of some which are normally solid. There are filters for removing particulate solids and sorption methods for removing I, and 3H,0 which have good potential for application to this off-gas, but no existing technology was found to remove and concentrate the Kr. The high concentration of CO, in the filtered gas stream ($^{\circ}90\%$ CO₂, 10% light gases, 10-20 ppm Kr) and the simílarity of CO, behavior to that of Kr preclude the use of commonly employed processes such as membrane permeation, sorption on charcoal or molecular sieves, and physical absorption by a third component. The removal of CO2 before such an operation appears feasible, but undesirable. A process involving Kr Absorption in Liquid CO2 (KALC), which exploits the fact that Kr is more soluble in liquid CO, than is O, N2, or CO, appears to be a natural solution to this particular problem.

The solubility of Kr in liquid ${\rm CO}_2$ was experimentally determined by an <u>in situ</u> radioactive counting method using ${\rm 85}{\rm Kr}$. This method avoided the problems usually associated with sampling systems of highly volatile components. Krypton tracer and ${\rm CO}_2$ were contained in a sealed stainless steel tube, brought to equilibrium by rocking in a constant-temperature bath, and counted with the tube in a vertical position to measure the radioactivity in each phase. Data were obtained at various temperatures over the entire ${\rm CO}_2$ liquid range.

Engineering calculations, based on literature values for the solubilities of O_2 , N_2 , and CO and the measured values for Kr, were made for a KALC system which employs a fractionator, a stripper, and an absorber. These calculations indicate that 99.9% of the Kr in the off-gas stream can be concentrated into a very small volume.

The Rare Gas Removal Pilot Plant at the ORGDP was used to explore the feasibility of operating a KALC system. In addition to demonstrating feasibility, the initial pilot campaign provided encouragement that the KALC process is a good way to decontaminate the HTGR reprocessing off-gases.

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Introduction

The off-gas from the reprocessing of HTGR fuel elements is unique among processing plant off-gases because the graphite and other carbon components of the HTGR fuel must be burned. The quantity of gas is larger than that evolved from LWR or LMFBR processing and the composition is different since it is mostly CO. Although alternative processing schemes which propose the separation of the graphite by physical methods to make burning unnecessary have been suggested, the problems encountered (not the least of which was the storage of the contaminated graphite) appear more difficult than treatment of the burner off-gas. The burning operations are performed at temperatures between 700 and 1000°C with a slight excess of oxygen to avoid, insofar as possible, the formation of CO. Commercial oxygen would be used rather than air to avoid handling large quantities of N_2 , but some inleakage of N, is unavoidable. It has been found (1) that several of the normally solid fission products are present in the off-gas at combustion conditions. The most important of the ones reported are Cs, Zr, Mb, Ru, and Ce, of which only the Ru exceeded (by a small amount) 10% volatilization. A sintered metal filter, located above the burner, was very effective in retaining all of these except Cs. About 2% of the Cs passed through a filter train, while the other metal fission products were retained with a decontamination factor greater than 103. The 3H, Xe, and Kr were all quantitatively released in the burning operations. Data on I, are not available, but it is also expected to be released during burning. An estimated analysis of the gas stream after filtration is given in Table 1. For a 1-metric ton/day plant, the given numbers would be multiplied by 100 since 100 blocks would be processed per day.

Since the removal and handling of Kr seemed to be the most important problem, our attention has been directed to it. The available technology was thoroughly assessed to determine whether a process that would effectively accomplish this task existed. (2) However, we found that the processes which are practical for removing Kr from streams similar to contaminated air, without seriously altering the composition of the bulk stream, are ineffective for streams containing large amounts of CO2. Indeed, the insidious similarity of the behavior of Kr and CO2 usually establishes tolerable limits of CO2 concentration in such processes. Sorption on molecular sieves or other solid sorbents, selective absorption by halocarbons, and selective membrane permeation techniques are among the processes eliminated by this factor.

The expedient of removing the CO₂ from the gas stream was then considered. This could be done by several methods. Reaction of the CO₂ with a lime bed produces prodigious amounts of solid radioactive waste for storage, and was thus ruled out. Caustic scrubbers are effective but also produce unacceptable amounts of waste. Among the regenerable scrubbing systems, the hot potassium carbonate system seems most desirable; specifically, it is as effective as, and more economical than, the amine sorbents. Finally, the possibility exists that the CO₂ could be removed by condensation. Because the triple point of CO₂ is -56.6°C at a vapor pressure of 5.1 atm, it is not feasible to completely remove the CO₂ as a liquid. In order to reduce the CO₂ content to a small fraction of the gas stream, desublimation temperatures of the order of -100°C are necessary even when the gas is compressed to a pressure of several atmospheres. None of the methods considered could ensure that the CO₂ removed from the gas stream would be sufficiently decontaminated from Kr or other fission products to allow its discharge. Further, the removal of CO₂ would not reduce the volume of the off-gas stream sufficiently to allow it to be stored, but would only prepare it for Kr removal by other methods.

Table 1. Estimated Burner Off-Gas Analysisa

Major Component	lb	scf	Mole Fraction
co ₂	1223	9976	0.897
02	75	846	0.075
N ₂	24	312	0.028
Contaminant	ppm (total)	ppm (radioactive)	Curies
Krypton	16	1.2	572.0
Tritium	2.4×10^{-2}	2.4×10^{-2}	9. 8
Iodine	3	9.2×10^{-7}	0.2
Xenon	61		

^aFor head-end reprocessing of one FSVR fuel block; 6-year exposure, 150-day cooling.

This rather unpromising review of existing technology led us into the development of a rather unique method for removing Kr from the HTGR burner off-gas. This method, which involves the selective absorption of Kr in liquid CO₂, has been called the KALC (Krypton Absorption in Liquid Carbon Dioxide) process. The process operates at about 20-atm pressure and at temperatures from about -45°C to -20°C. The flowsheet, in simplified form, is shown in Fig. 1. The flow rates shown are those that might be used in a pilot facility and would be equivalent to processing three blocks of Ft. St. Vrain fuel per day. The flow ratios shown are approximate; however, more precise calculations, which continually change with incorporation of newly acquired data, show no qualitative differences.

Before being fed into the system, the filtered burner off-gas is compressed and chilled so that part of the CO₂ is condensed. In the absorber-fractionator system, virtually all of the Kr is selectively absorbed in the liquid phase, while all gas components with solubilities less than Kr (e.g., O₂, N₂ and CO) are returned to the gas phase and discharged from the top of the absorber along with much of the CO₂. Gases more soluble than Kr (e.g., Xe) follow the liquid CO₂ to the stripper, where the Kr is removed and concentrated. The CO₂ from the bottom of the stripper would contain components of low volatility. Xenon either could be allowed to accumulate in the CO₂ to the point where it is eventually discharged from the top of the absorber, or it could be removed by a separate distillation operation on the recycle CO₂ stream. This technique should yield a xenon product of very high purity.

The scrubber may be expected to remove from the incoming gas most of the particulates that reach the process. Under KALC conditions, water and all conceivable forms of iodine are essentially nonvolatile components, which, at the expected inlet concentrations, are soluble in the liquid phase. Solids and nonvolatile components would tend to accumulate in the recycled liquid CO_2 . They could be removed by an evaporator, which would return purified CO_2 to the process and concentrate the solids for discharge. When it is realized that the volatilities of water and the iodine compounds are limited by the vapor pressure of the pure solid phases, one can imagine such an evaporator effectively operating with a thick slurry of crystals in the reboiler.

Development of the KALC process requires: (1) obtaining the necessary equilibrium and thermodynamic data for describing the system; (2) selecting, characterizing, and, in some cases, developing components required for a radio-chemical application of the process; (3) selecting an effective operating mode with its associated control system; and (4) demonstrating the performance of an integrated system. At this time, items (2) and (3) have received only sufficient attention to assure us that appropriate solutions will be forthcoming. Work on items (1) and (4) has already yielded significant results.

Solubility of Krypton in Liquid CO2

An experimental determination of the solubility of krypton in liquid CO $_2$ had been made at KFA Jülich previously, but the results differed considerably from the solubility calculated by assuming ideal behavior. Therefore, we made our own measurements using in situ counting of tracer 85 Kr distributed between equilibrated gaseous and liquid phases in a sealed container. The in situ technique eliminated the inaccuracies usually associated with sampling highly volatile systems.

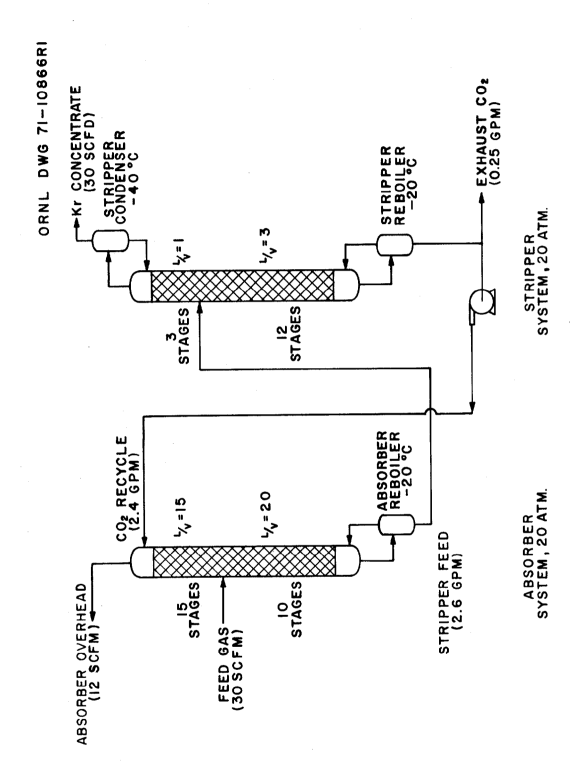


Fig. 1. Typical Flowsheet for the KALC Process.

The equipment used is shown schematically in Fig. 2. Essential components were a cold bath, an equilibrating device, and counting equipment. The temperature of the cold bath was controlled to within $\pm 0.03^{\circ}\text{C}$ by circulating a slight excess of coolant from a dry ice--trichlorethylene reservoir and supplying heat as called for by a thermostat. The Kr-CO₂ mixture was contained in a l-in.-diam by ll-in.-long, stainless steel cylinder, which was rocked for at least 1 hr at temperature. We have determined that distribution equilibrium is attained by this procedure. The equilibrated system was then set upright, and the ^{85}Kr radioactivity was counted in each phase, yielding raw data on a volume basis. Data reduction involved correcting for internal attenuation and backscatter from the CO₂, and recalculating the Kr activity on a mole (rather than volume) basis. The correction, which was sometimes as much as 25% (depending on the CO₂ density and the gamma energy range counted), was made by means of a calibration curve. This curve was prepared empirically by incrementally adding CO₂ to a fixed quantity of Kr. The recalculation merely required the use of the CO₂ density, which is well known.

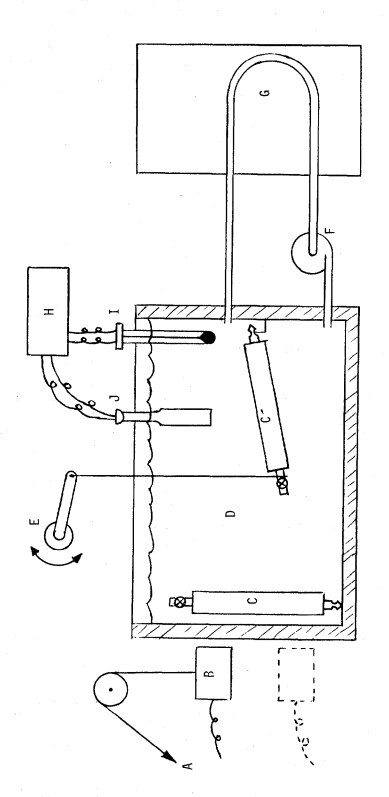
The data were finally expressed as a separation factor, Y_{Kr}/X_{Kr} , where Y and X are the mole fractions of Kr in the gaseous and liquid phases, respectively. The total Kr present (85 Kr plus nonradioactive Kr) was usually less than 1000 ppm; however, since the separation factor is a ratio, it was not necessary to know the exact amount. The reduced data are shown in Fig. 3. Fifty-nine data points, covering the entire liquid range, were obtained. The curve drawn through our data is linear over most of its length, showing a slight positive curvature near the triple point and dropping off strongly toward a value of unity as the critical temperature is approached. Our curve has about twice the slope of the calculated ideal curve and crosses it near 0°C. A more sophisticated calculated curve, which allows for nonideality in both phases, (4) falls below the ideal curve. The Jülich data tend to follow the latter curve but show a minimum at about 20°C. The fact that no minimum was predicted favors our results. Further, in order to validate the method used here, the distribution of 85 Kr in a Kr-H₂O system was determined and found to agree with reported values obtained by other methods. (5) Our known errors (from uncertainties in counting, temperature, CO₂ density, and empirical calibration) are somewhat less than the total scatter shown in Fig. 3.

Preliminary data indicate that the distribution of Kr is not significantly altered either at much higher Kr contents or by the presence of added oxygen.

Solubility of O2, N2, and CO in CO2

The literature has provided a reasonable amount of data on the solubility of O_2 in liquid CO_2 , (6-8) somewhat less on N_2 in CO_2 , (6,8,9) and a very small amount on CO in CO_2 . Most of the reported data were obtained at high concentrations, and hence at higher pressures than are of interest to this work. When the light gases comprise less than 50% of the gas phase, at total pressures less than 50 atm, the liquid phase contains less than 10% dissolved light gases. As the concentration of light gases decreases from this level, data become sparse and the accuracy diminishes; however, theoretical considerations have helped us to extrapolate the data into the low-concentration regions.

Equilibrium data from the literature have been fitted to a model which, over the range of conditions of interest, describes the KALC system quite satisfactorily. Of necessity, the model is based on binary system data. The only ternary data



F - Circulating bump G - Dry ice/solvent cold reservoir H - Temnerature controller I - Sensing element J - 250 watt heater

CounterweightLead-shielded scintillation counter

Cylinder in counting position Cylinder in equilibrating position Controlled temperature bath

A B O O D B B

Motor-driven oscillator

Fig. 2. Apparatus for in situ Measurement of Equilibrium Distribution of $^{85}\mathrm{Kr}$ Between CO_2 Liquid and Gas Phases.

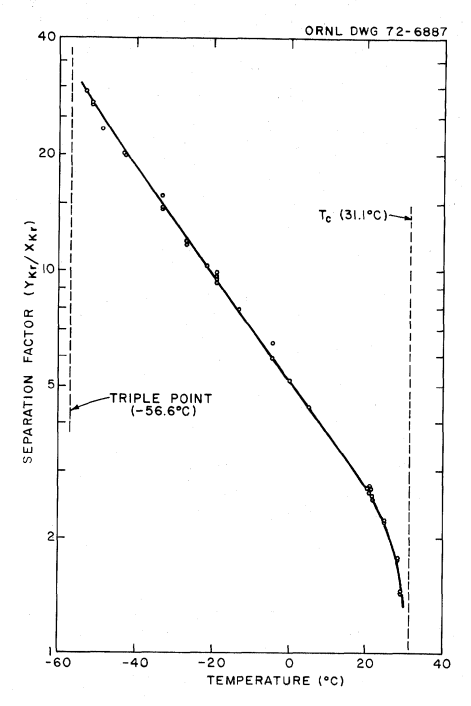


Fig. 3. Distribution of Krypton Between Gaseous and Liquid CO2.

obtained under conditions close to those used for the KALC system were for the $\mathrm{O_2-N_2-CO_2}$ system at pressures higher than 50 atm. Even though they were out of the proper range, these data and equilibrium numbers calculated from the model were in reasonable agreement. The model is most simply expressed in the following form: given a liquid phase composition and a temperature, calculate the total pressure and the vapor phase composition.

The total pressure was found to fit the empirical relation

$$P_{t} = P_{CO_{2}}^{*} + \sum_{i} x_{i} + \sum_{i} x_{i} + \sum_{i} x_{i} + \sum_{i} x_{i}$$
(1)

where

 $P_{+} = total pressure (atm),$

 $P_{CO_2}^*$ = saturated vapor pressure of CO_2 at system temperature (atm),

x = mole fraction in liquid phase,

i,j = index subscripts applied to all components except CO2,

a,b,c = empirically determined constants.

The compressibility of CO₂ vapor is such that a correction for nonideality is required. All other components have critical points sufficiently removed from KALC conditions that gas phase ideality can be assumed. A reasonable form for the gas phase composition then becomes:

$$y_{CO_2} = (p_{CO_2}/z)/[\Sigma p_j + (p_{CO_2}/z)]$$
 (2)

$$y_i = p_i / [\Sigma p_j + (p_{CO_2}/\Sigma)],$$
 (3)

where

Z =the compressibility factor for saturated CO_2 at the temperature of the system,

p = partial pressure,

y = mole fraction in the vapor phase.

The partial pressures are defined so that

$$P_{T} = P_{CO_{2}} + \Sigma P_{i}. \tag{4}$$

Because the concentration of the dissolved light gases is small, Raoult's law was applied to the CO₂. It was necessary, however, to make a Poynting-type correction of the CO₂ vapor pressure. As an expedient in getting a better fit, the molar volume of the liquid phase was not used; instead, a fictitious molar volume about a factor of 3 times the real molar volume was employed.

The partial pressure of CO2 is, then,

$$P_{CO_2} = P_{CO_2} (1 - \Sigma x_j), \qquad (5)$$

where

$$P_{CO_2} = P_{CO_2}^* \exp[k(P_t - P_{CO_2}^*)/T],$$
 (6)

and

 P_{CO_2} = effective vapor pressure of CO_2 (atm),

T = absolute temperature (°K),

k = a determined constant = 1.7 °K/atm.

The partial pressure of the ith component is given by:

$$P_{i} = \frac{x_{i}}{\sum x_{j}} [P_{CO_{2}}^{*} - P_{CO_{2}}^{*} (1 - \sum x_{j})] + a_{i}x_{i} + b_{i}x_{i}P_{CO_{2}}^{*} + c_{i}x_{i}\sum x_{j}.$$
 (7)

For a binary system, Equation 7 reduces to:

$$P_i = P_t - (1 - x_i)P_{CO_2}.$$
 (8)

These equations comprise a set from which equilibrium can be calculated. Table 2 gives the numerical values for the constants, along with the expressions used in this study to compute the compressibility factor, Z, and the saturated vapor pressure of CO_2 , P_{CO_2} .

Henry's law constants consistent with the model, and defined as the limit of the ratio of the partial pressure to the mole fraction in the liquid phase, are shown in Fig. 4. It is noted that CO is predicted to be the most difficult component to separate from Kr, but only slightly more difficult than oxygen. The Kr-CO separation factor increases from 1.43 at 0°C to 1.8 at -20°C. These values are sufficiently high that neither the required height of towers nor the control of flow rates for KALC will cause undue concern.

Engineering Feasibility of the KALC Process

Despite the early stage of the development of KALC, the feasibility of the process has been demonstrated in a recent campaign conducted by ORNL at the ORGDP Rare Gas Removal Pilot Plant. The primary objective of this campaign was to establish our confidence in operating a liquid-CO2--light-gas system. However, ⁸⁵Kr was used in tracer amounts during much of the operation.

The basic pilot plant, which has been used quite extensively over the past few years in the development of the fluorocarbon absorption processes, is composed of three separate columns for carrying out the absorption, fractionation, and stripping operations. Each of the packed columns has a 9-ft section of packing; the absorber and the fractionator are each 3 in. in diameter, and the stripper is 10 in. in diameter.

A nominal gas flow of 10 to 15 scfm to the absorber, with a countercurrent flow of about 1.5 gpm of liquid ${\rm CO}_2$, was used during the feasibility studies. Under these conditions and with an operating pressure of about 20 atm, the column temperatures were in the predicted range of -20 to -40°C. The system operated very smoothly throughout the campaign period (three weeks).

In-line analyses were not possible for checking concentrations, etc., but estimates (which were later verified by mass spectrographic analyses) leave little

Coefficients Used to Correlate Equilibrium Data for the KALC System Table 2.

Component i	В	q	υ
02	532.33	-2.7153	-494.72
$^{ m N}_2$	720.22	-5.9729	-553.88
000	484.39	-1.8473	-395.10
Kr	188.0	2.80	-

Total Pressure = $P_t = P_{CO_2}^* + a_1x_1 + b_1x_1P_{CO_2}^* + c_1x_1^{1/3}$

Henry's Constant = $H_i = a_i + P_{CO_2}^* (1 + b_i)$

Saturation Vapor Pressure of ${\rm CO}_2$:

ln (P,*) = -8.175626 + 26.27742
$$\left(\frac{T}{600}\right)$$
 - 23.76839 ln $\left(\frac{T}{600}\right)$ - 8.62709785 $\left(\frac{600}{T}\right)$

Compressibility of CO $_2$ = Z = 3.21203 - 0.029634967T + 1.3757 x $_10^{-4}$ T $_2$ - 2.2960 x $_10^{-7}$

 $T = Absolute temperature ({}^{\circ}K)$



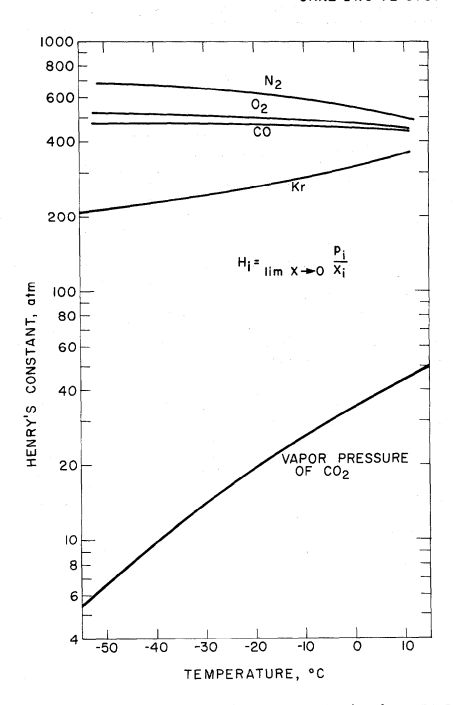


Fig. 4. Henry's Constants for Major Components in the KALC Process.

doubt that such a system, with controls and instrumentation suited specifically to the liquid CO₂ process, can effectively remove Kr from the burner-type off-gases. Spot checks on the behavior of Kr in the system at arbitrary run conditions indicated both absorber and process DFs on the order of 100, with process concentration factors in excess of 1000.

Discussion

An economic evaluation of the KALC process has not yet been undertaken; thus specific costs are not yet available. The compressor and refrigeration requirements probably account for a major portion of the total expense for the KALC system, but even these costs will not be severe. The capital and operating costs associated with the KALC system should be only a small fraction of the costs for the rest of the reprocessing plant. Corrosion in the basic KALC process should present no problems. However, more study is required on corrosion in any special operation, such as one that would concentrate iodine and tritium for removal.

Many questions of an engineering nature are still unanswered; primarily, these questions are concerned with equipment design and performance, but they also include the selection of the operating mode most conducive to reliable control and system performance under transient loads. Preliminary analysis of the control problem indicated several plausible alternatives. A calculational model of the KALC process which can predict system performance will be valuable both in studying control and in interpreting the results of engineering experiments. Such a model is now partially complete.

Doubts about the feasibility of the KALC process have been dispelled by an actual demonstration in engineering equipment and by the accumulation and consideration of basic equilibrium and physical property data. We have yet to discover any supportable objection to the application of this process to HTGR off-gas decontamination. Our determination of the distribution of Kr in the KALC system and/or the data on the other light gases reported in the literature would have to be in error by at least a factor of 2 (which we consider incredible) to render the process unworkable. In summary, then, the advantages of using one of the components of the stream as the solvent to effect its decontamination, the moderate operating conditions employed, and the general effectiveness of the process make KALC very attractive.

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DISCUSSION

LASER: In our lab, we have a poster which displays: "Atomic Power means clean air." That's right. But we all know that we must add: Reprocessing means emission of krypton, tritium, and possibly 129 iodine.

A special problem of reprocessing is the decontamination of the burner off-gas of THTR fuel. This is nearly the same as HTGR fuel, which you have heard of. Therefore, since 1970, we have made theoretical studies and laboratory experiments of suitable processes to retain volatile and non-volatile activity. During the last year we have made several hot cell experiments and have installed a complete decontamination process for hot cell experiments called AKUT.

Let me give you a short description of the results and of the process. The composition of the burner off-gas differs considerably depending on experimental conditions. The carbon monoxide content fluctuates between approximately 5 and 25%. The activity, especially of 85 krypton, fluctuates by a factor of 10 or more during a short time because of irregular burning of the coated particles. Therefore, we have averaged measurements over an extended period to arrive at mean values. In the case of carbide fuel elements the burner off-gas activity is approximately 20 mCi/m³ tritium, 420 mCi/m³ krypton, and nearly 125 mCi/m³ cesium. In the case of oxide fuel elements, the emitted activity is lower; approximately by a factor of 10.

Most of the off-gas activity is associated with aerosols, especially cesium and iodine when short cooled fuel elements are burnt. More than 99.8% of the cesium activity is separable by an electrostatic precipitator. The iodine activity decontamination factor has fluctuated between 2 and 2,000 because of different ratios of particulate to molecular species. With absolute filters downstream of the electrostatic separator, cesium activity can be reduced below the maximum permitted concentration. The iodine activity, however, remains high by a factor up to 1,000.

We have developed a process to separate krypton from the off-gases by scrubbing with liquid carbon dioxide. We achieved decontamination factors of 600 and more. In practice, because we have a gas with fluctuating composition, control may be difficult. A further problem is enrichment of krypton for storage and disposal. We now prefer to liquify all of the off-gas by pressurizing at room temperature and then, to enrich the krypton by rectification.

The hot cell experiments consist of the following steps:

- (1) Precipitation of aerosols by an electrostatic separator.
- (2) Decontamination of off-gases by an absolute filter. (We prefer a relatively clean off-gas before going into krypton separation.)

- (3) Oxidation of carbon monoxide with oxygen to give an off-gas with approximately 95% carbon dioxide and 5% oxygen.
- (4) Liquifying the off-gas by compression to approximately 70 atmospheres.
- (5) Enriching the krypton by rectification to approximately 10 to 20% krypton.
- (6) Further rectification to enrich tritiated water and, possibly, iodine at the bottom of the rectification column.
- (7) Evaporation of the tritium and iodine enriched solution and passing the gas over a tritium and iodine absorber.

A very coarse estimation of cost is 1.5×10^{-5} DM per KWH, but this value can be higher by a factor of 2.